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***Neofomitella polyzonata* gen. et sp. nov., and
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ABSTRACT — Phylogenetic analysis based on ITS, nLSU, and RPB2 sequences revealed that *Fomitella* in the current sense belonged to two distantly related subclades in the core polyporoid clade. *Fomitella* in a narrow sense is proposed for the type species, *F. supina*, and *Neofomitella* gen. nov. is proposed for *Fomitella fumosipora* and *F. rhodophaea* and a new species *N. polyzonata*. *Neofomitella* differs from *Fomitella* by its distinctly crusted basidiocarps with the cuticle developing from base to margin. Illustrated descriptions of the new genus and species are provided. The main morphological differences between *Fomitella*, *Neofomitella*, and related genera are discussed, and an identification key to *Neofomitella* is also provided.

KEY WORDS — phylogeny, *Polyporaceae*, *Polyporales*, *Basidiomycota*, taxonomy

Introduction

Fomitella Murrill, typified by *F. supina* (Sw.) Murrill, was erected as a monotypic genus by Murrill (1905). Subsequently, *F. fumosoavellanea* (Romell) Murrill [*Trichaptum fumosoavellaneum* (Romell) Rajchenb. & Bianchin.], *F. fumosipora*, *F. rhodophaea*, and *F. malaysiana* (Corner) T. Hatt. & Sotome were transferred to the genus (Murrill 1908; Hattori 2005; Hattori & Sotome 2013). Hattori (2005) emended the generic concept of *Fomitella* as follows: basidiocarps annual to perennial, effused-reflexed to distinctly pileate, pileal surface glabrous to minutely tomentose, context firm-fibrous to corky, light orange to pale brown, with a dark agglutinated crust; hyphal system trimitic with clamped generative hyphae, skeletal and binding hyphae well differentiated, negative in Melzer's reagent; cystidia absent; basidiospores

ellipsoid to cylindrical, colorless, thin-walled, smooth, and negative in Melzer's reagent; causing a white rot.

On-going studies on the diversity of wood-rotting fungi in eastern China have produced several new species (Dai & Cui 2005; Cui & Dai 2008a,b; Cui et al. 2007, 2008; He & Dai 2012; Wang et al. 2009, 2011; Cao et al. 2012), and additional study on the specimens from this area recently revealed an undescribed species. Phylogenetic analysis clustered our undescribed species with *Fomitella fumosipora* and *F. rhodophaea* as a clade separated from *F. supina*. Therefore, we propose a new genus *Neofomitella* to accommodate the new species and two new combinations from *Fomitella*.

Materials & methods

Morphological studies

The studied specimens were deposited at the herbaria of the Institute of Microbiology, Beijing Forestry University (BJFC) and Institute of Applied Ecology, Chinese Academy of Sciences (IFP). The microscopic procedure follows Cui & Zhao (2012). To present the basidiospore size variation, 5% of measurements were excluded from each end of the range and were given in parentheses. In the text the following abbreviations were used: IKI = Melzer's reagent, IKI- = both inamyloid and non-dextrinoid, KOH = 5% potassium hydroxide, CB = Cotton Blue, CB- = acyanophilous, L = mean spore length (arithmetic average of all spores), W = mean spore width (arithmetic average of all spores), Q = variation in the L/W ratios between the specimens studied, n = number of spores measured from given number of specimens. Special colors follow Petersen (1996).

Molecular study and phylogenetic analysis

Dried specimens were used for molecular study. CTAB rapid Plant genome extraction kit-DN14 (Aidlab Biotechnologies Co. Ltd, Beijing) was used to extract DNA, according to the manufacturer's instructions with some modifications. ITS region was amplified with primer pair ITS5 and ITS4 (White et al. 1990), nLSU region with primer pair LR0R and LR7 (<http://www.biology.duke.edu/fungi/mycolab/primers.htm>), and RPB2 with primer pair rRPB2-f5F and bRPB2-7.1R (Liu et al. 1999; Matheny 2005). The PCR procedures were: (1) for ITS — initial denaturation at 95°C for 3 min, followed by 35 cycles at 94°C for 40 s, 54°C for 45 s and 72°C for 1 min, and a final extension of 72°C for 10 min; (2) for nLSU — initial denaturation at 94°C for 1 min, followed by 35 cycles at 94°C for 30 s, 50°C for 1 min and 72°C for 1.5 min, and a final extension of 72°C for 10 min; and (3) for RPB2 — initial denaturation at 94°C for 2 min, followed by 10 cycles at 94°C for 40 s, 60°C for 40 s and 72°C for 2 min, then followed by 37 cycles at 94°C for 45 s, 55°C for 1.5 min and 72°C for 2 min, and a final extension of 72°C for 10 min. The PCR products were directly sequenced in Beijing Genomics Institute, China, with the same primers.

Other reference sequences for our phylogenetic analysis were selected from Binder et al. (2005), Justo & Hibbett (2011), Miettinen & Rajchenberg (2012), and BLAST searches in GenBank (TABLE 1). These sequences were sampled from *Polyporales* with

TABLE 1. Polyporalean and outgroup taxa used in phylogenetic analysis.
(Sequences generated in this study are in bold.)

SPECIES	SAMPLE NUMBER	ITS	nLSU	RPB ₂
<i>Abundisporus pubertatis</i>	Dai 11927	KC867398	KC867494	KF274654
<i>Abundisporus violaceus</i>	MUCL38617	FJ411100	FJ393867	—
<i>Climacodon septentrionalis</i>	AFTOL-ID 767	AY854082	AY684165	AY780941
<i>Corioloopsis aspera</i>	Cui 6702	KC867353	KC867476	KF274658
	Cui 6725	KC867356	KC867477	KF274659
<i>Corioloopsis brunneoleuca</i>	Dai 12087	KC867416	KC867435	KF274656
	Dai 12180	KC867414	KC867432	KF274655
<i>Corioloopsis byrsina</i>	FP-105050-Sp	JN165001	JN164788	JN164871
<i>Corioloopsis cf. caperata</i>	CR22	JN164999	JN164789	JN164870
	Ryvarden 45481	KC867399	KC867428	KF274657
<i>Corioloopsis retropicta</i>	Dai 9870	KC867404	KC867443	KF274653
<i>Corioloopsis rigida</i>	BJFC12680	KC867381	KC867454	KF274664
<i>Corioloopsis sanguinaria</i>	Cui 5444	KC867387	KC867463	—
	Dai 9314	KC867390	KC867467	—
<i>Corioloopsis sp.</i>	BRFM1125	JX082370	—	—
	BRFM1126	JX082371	—	—
<i>Corioloopsis strumosa</i>	Dai 10642	JX559278	JX559303	JX559312
	Dai 10657	KC867371	KC867491	KF274650
<i>Daedaleopsis confragosa</i>	Cui 9732	JX569731	JX569748	KF274647
<i>Daedaleopsis sinensis</i>	Dai 11431	JX569732	JX569749	KF274648
<i>Datronia mollis</i>	RLG6304sp	JN165002	JN164791	JN164872
<i>Datronia scutellata</i>	RLG9584T	JN165004	JN164792	JN164873
<i>Dentocorticium sulphurellum</i>	T609	JN165015	JN164815	JN164875
<i>Donkioporia expansa</i>	P188	HM536087	HM536052	HM536102
<i>Earliella scabrosa</i>	PR1209	JN165009	JN164793	JN164866
<i>Fomes fomentarius</i>	Cui 8020	JX290073	JX290070	—
<i>Fomitella supina</i>	JV0610	KF274645	KF274646	—
	Nunez 1183	KF274644	—	—
	Ryvarden 39027	KF274643	—	—
<i>Fomitopsis pinicola</i>	AFTOL-ID 770	AY854083	AY684164	AY786056
<i>Funalia gallica</i>	BJFC12697	KC867379	KC867453	—
	RLG-7630-sp	JN165013	JN164814	JN164869
<i>Funalia trogii</i>	RLG-4286-Sp	JN164993	JN164808	JN164867
<i>Ganoderma tsugae</i>	AFTOL-ID 771	DQ206985	AY684163	DQ408116
<i>Grifolia sordulenta</i>	AFTOL-ID 562	AY854085	AY645050	AY786058
<i>Hexagonia apiaria</i>	Cui 6447	KC867362	KC867481	KF274660
<i>Hexagonia glabra</i>	Cui 8468	JX559277	JX559302	JX559311
	Dai 10991	JX569733	JX569750	KF274649
<i>Lignosus rhinocerotis</i>	PEN94	JQ409359	AB368074	AB368132
<i>Lopharia cinerascens</i>	FP-105043-sp	JN165019	JN164813	JN164874
<i>Megasperia major</i>	Cui 10253	JQ314366	JQ780437	JX559314
<i>Megasporoporiella subcavernulosa</i>	Cui 9252	JQ780378	JQ78041	JX559315
<i>Microporus affinis</i>	Cui 7714	JX569739	JX569746	KF274661
<i>Microporus flabelliformis</i>	Dai 11574	JX569740	JX569747	KF274662
<i>Microporus xanthopus</i>	Cui 8284	JX290074	JX290071	JX559313
<i>Neofomitella fumosipora</i>	Cui 8816	JX569734	JX569741	—
	Dai 10777	JX569735	JX569742	—
<i>Neofomitella polyzonata</i>	Dai 10419	JX569738	JX569745	KF274663
	Dai 10420	JX569736	JX569743	—
	Dai 11360	JX569737	JX569744	—
<i>Neofomitella rhodophaea</i>	TFRI 414	EU232216	EU232300	—
<i>Perenniporia corticola</i>	Cui 1465	JN048759	JN048779	KF274651
<i>Perenniporia tenuis</i>	Wei 2783	JQ001858	JQ001848	KF274652
<i>Phlebia radiata</i>	FPL6140	AY854087	AE287885	AY218502
<i>Polyporus granmocephalus</i>	WD2343	AB587626	AB368089	AB368146
<i>Polyporus varius</i>	WD2347	AB587636	AB368111	AB368168
<i>Pseudofavolus cucullatus</i>	WD2157	AB587637	AB368114	AB368170
<i>Trametes betulina</i>	HHB-9942-sp	JN164983	JN164794	JN164860
<i>Trametes elegans</i>	FP-105679-sp	JN164944	JN164799	JN164861
<i>Trametes polyzona</i>	BKW-004	JN164978	JN164790	JN164856
<i>Trametes sanguinea</i>	PR-SC-95	JN164982	JN164795	JN164858
<i>Trametes suaveolens</i>	FP-102529-sp	JN164966	JN164807	JN164853
<i>Trametes versicolor</i>	FP-135156-sp	JN164919	JN164809	JN164850
<i>Trametopsis cervina</i>	TJV-93-216-sp	JN165020	JN164796	JN164877
<i>Boletopsis leucomelaena</i>	AFTOL-ID 1527	DQ484064	DQ154112	GU187820
<i>Hydnellum geogenum</i>	AFTOL-ID 680	DQ218304	AY631900	DQ408133

the aim of providing representatives for all major clades within the order, concluded by Larsson et al. (2004), Binder et al. (2005) and Larsson (2007). *Boletopsis leucomelaena* (Pers.) Fayod and *Hydnellum geogenium* (Fr.) Banker were selected as outgroup (Justo & Hibbett 2011). Phylogenetic analysis for each single gene was carried out, and we got similar topologies to combined dataset in both MP and Bayes analyses. So in the paper we showed the result from the combined dataset that received higher support values. This combined dataset was aligned using Clustalx 1.83 (Chenna et al. 2003) and manually edited as necessary. Sequence alignment was deposited at TreeBase (submission ID 14418).

Maximum parsimony (MP) analysis was performed using PAUP* 4.0b10 (Swofford 2002) with gaps treated as missing data. Trees were generated using 100 replicates of random stepwise addition of sequence and tree-bisection reconnection (TBR) branch-swapping algorithm. All characters were given equal weight. Branch support for all parsimony analysis was estimated by performing 1 000 bootstrap (BP) replicates (Felsenstein 1985) with a heuristic search of 10 random-addition replicates for each bootstrap replicate.

The best-fit model of nucleotide substitution was selected by hierarchical likelihood ratio tests (hLRT, Huelsenbeck & Crandall 1997; Posada & Crandall 2001) implemented in the MrModelTest 2.2 (Posada & Crandall 1998; Nylander 2004). Following this model, MrBayes 3.1.2 (Ronquist & Huelsenbeck 2003) was used for Bayesian inference (BI). Eight Markov chains were run from random starting tree for 3,000,000 generations and sampled every 100 generations. The first one-fourth of the trees, which represented the burn-in phase of the analysis, were discarded, while the last three-fourths were used for calculating Bayesian posterior probabilities (BPPs) in the consensus tree. Confident branch support is defined as BPPs not less than 0.95 and bootstrap values above 50%.

Results

Molecular phylogeny

The combined dataset included 66 samples with 66 ITS, 62 nLSU, and 51 RPB2 sequences, representing 50 species and two collections not identified to species level from the main clades of *Polyporales*, as well as two species from *Thelephorales* as outgroup. MP analysis yielded eight equally parsimonious trees (tree length = 9 153, CI = 0.283, RI = 0.520, RC = 0.147, HI = 0.717). The best model for the alignment was estimated as "GTR+I+G". BI resulted in an average standard deviation of split frequencies 0.005127. The topologies from MP and BI were similar, and only the BI tree was presented along with the BP from MP analysis (FIG. 1). In the phylogenetic analysis, sampled species of *Fomitella* formed two subclades within the core polyporoid clade. One subclade comprised *Fomitella supina*, the generic type, and two unidentified collections (labelled as *Corioloropsis*) from French Guiana (BPP = 1.00, MP = 100%; FIG. 1), while the second subclade included *F. fumosipora*, *F. rhodophaea* and our undescribed species (BPP = 1.00, MP = 96%; FIG. 1). Taken the morphology into consideration together, we propose a new genus, *Neofomitella*, to accommodate

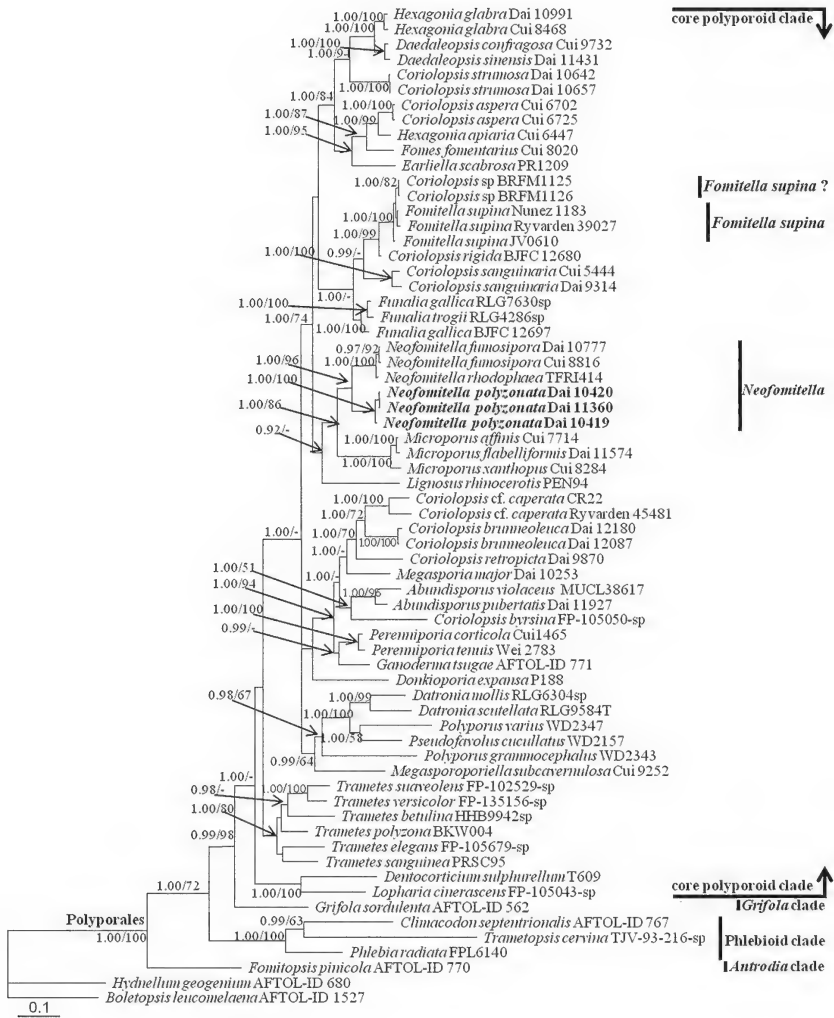


FIG. 1. Phylogram of polyporalean taxa obtained from Bayesian inference of the combined dataset of ITS, nLSU, and RPB2. Bayesian posterior probabilities more than 0.95 and bootstrap values above 50% are indicated above or below the branches. The scale bar represents estimated number of changes per site.

the second subclade; we transfer the two *Fomitella* species to the new genus and describe our new species as *N. polyzonata*. Phylogenetically, the *Neofomitella* subclade clusters with a *Microporus* subclade (FIG. 1).

Taxonomy

Neofomitella Y.C. Dai, Hai J. Li & Vlasák, **gen. nov.**

MYCOBANK MB 804799

Differs from *Fomitella* by its distinctly crusted basidiocarps with the cuticle developing from base to margin and from *Microporus* by its buff, yellowish brown, brown to pale grey context.

TYPE SPECIES — *Polyporus rhodophaeus* Lév.

ETYMOLOGY — *Neofomitella* (Lat.): referring to the morphological similarity to *Fomitella*.

Basidiocarps annual or perennial, pileate, sessile or effused-reflexed. Pileal surface yellowish-brown, brown, orange-brown, reddish-brown, fuscous to almost black, usually concentrically zonate or sulcate, glabrous to velutinate. Context buff, yellowish brown, brown to pale grey, corky to hard corky, with a dark agglutinated crust developing from base to margin. Pore surface usually white, cream to pale buff when fresh, pale brown to yellowish-brown when dry. Hyphal system trimitic with clamped generative hyphae, skeletal and binding hyphae well differentiated, negative in Melzer's reagent. Cystidia absent. Basidiospores oblong ellipsoid to cylindrical, hyaline, thin-walled, smooth, IKI–, CB–, tissue turn into black in KOH. Growing usually on angiosperm wood and causing a white rot.

Neofomitella fumosipora (Corner) Y.C. Dai, Hai J. Li & Vlasák, **comb. nov.** FIG. 2a

MYCOBANK MB 804803

≡ *Trametes fumosipora* Corner, Beih. Nova Hedwigia 97: 106 (1989).

≡ *Fomitella fumosipora* (Corner) T. Hatt., Mycoscience 46: 309 (2005).

SPECIMENS EXAMINED — CHINA. GUANGDONG PROVINCE, SHIXING COUNTY, Chebaling Nature Reserve, fallen angiosperm trunk, 25 Jun 2010, B.K. Cui 8816 (BJFC 7756); 23 Nov 2010, B.K. Cui 8715, 8717 (BJFC 7657, 7659); HAINAN PROVINCE, CHANGJIANG COUNTY, Bawangling Nature Reserve, fallen angiosperm trunk, 8 May 2009, Y.C. Dai 10777 (BJFC 5021). MALAYSIA. NEGERI SEMBILAN, Pasoh For. Res., 5 Dec 1998, T. Hattori (BJFC: ex. TFM ex. F. 19017); PENANG, Penang Hill, alt. 500–750 m, 13 Dec 2002, T. Hattori (BJFC: ex. TFM ex. F. 20477).

Neofomitella polyzonata Y.C. Dai, Hai J. Li & Vlasák, **sp. nov.**

FIGS 2b,c, 3

MYCOBANK MB 804804

Differs from *Neofomitella fumosipora* and *N. rhodophaea* by its distinctly velutinate pileal surface and larger pores.

TYPE — China, Jiangxi Province, Fenyi County, Dagang Mountain, on fallen trunk of *Cyclobalanopsis blakei* (Skan) Schottky (Fagaceae), 18 Sep 2008, Y.C. Dai 10419 (holotype, BJFC 4668).

ETYMOLOGY — *polyzonata* (Lat.): referring to the multiple zones on the pileal surface.

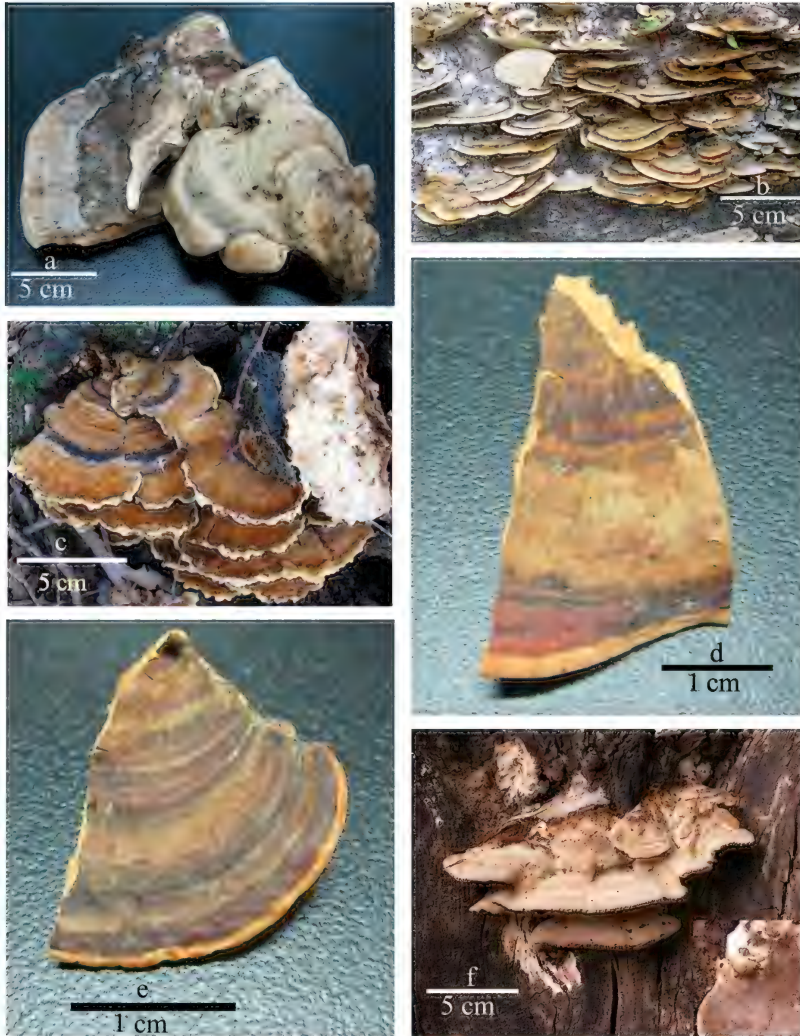


FIG. 2. *Neofomitella* and *Fomitella* basidiocarps. a: *N. fumosipora*. b, c: *N. polyzonata*. d, e: *N. rhodophaea*. f: *F. supina*.

FRUITBODY — Basidiocarps annual, pileate, sessile, usually imbricate, without odor or taste when fresh, hard corky to woody hard and light in weight upon drying. Pilei applanate, semicircular to dimidiate, up to 6 cm long, 10 cm wide and 6 mm thick at base. Pileal surface buff-yellow, curry-yellow, cinnamon, orange-brown to reddish brown, with one or more vinaceous brown, dark blue

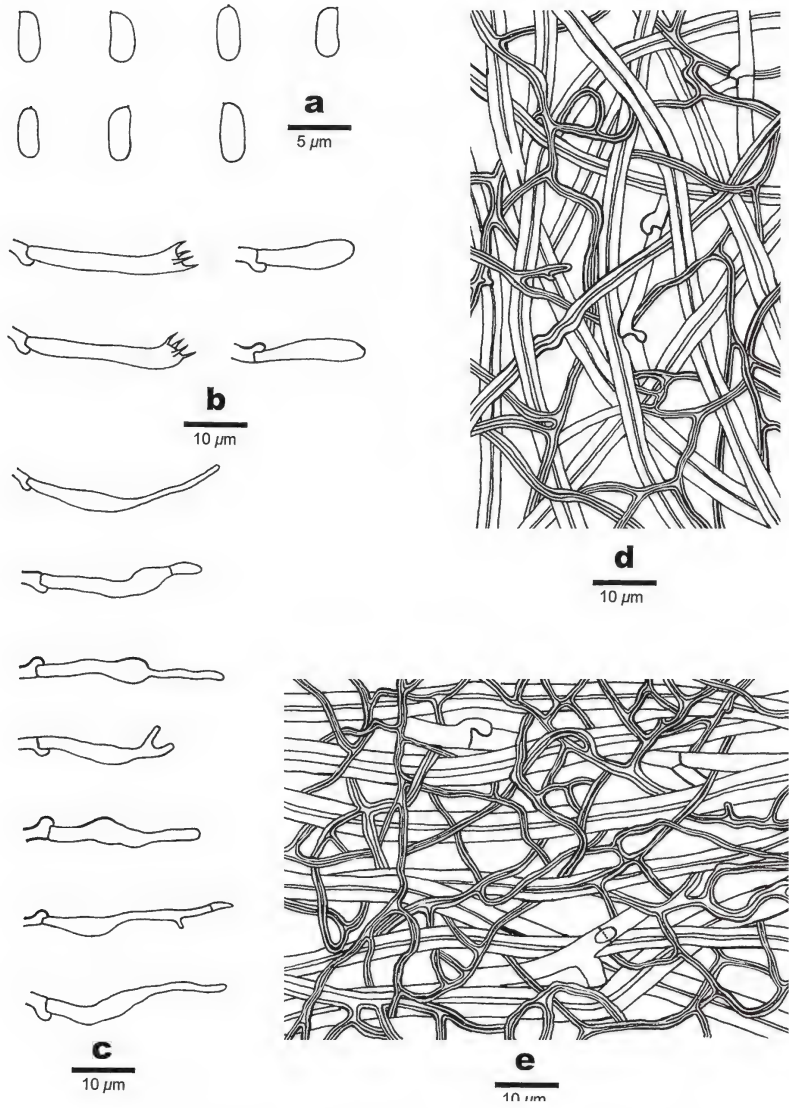


FIG. 3. *Neofomitella polyzonata* (drawn from the holotype).
a: Basidiospores. b: Basidia and basidioles. c: Cystidioles.
d: Hyphae from trama. e: Hyphae from context.

to almost black zones, finely velutinate, concentrically zonate; white to cream outgrowth occasionally spreading from the base with age. Margin cream, buff to buff-yellow, usually acute or slightly wavy. Pore surface cream to buff when

fresh, buff to yellowish brown when dry or bruised; sterile margin indistinct, white to cream, up to 0.5 mm wide; pores round, 3–4 per mm; dissepiments thin, entire. Context buff to yellowish brown, hard corky, azonate, up to 3 mm thick, a more or less dark agglutinated crust present as black zones in context towards upper surface. Tube layer concolorous with pore surface, up to 3 mm long.

HYPHAL STRUCTURE — Hyphal system trimitic; generative hyphae bearing clamp connections; skeletal and binding hyphae IKI–, CB–; tissue turning into black in KOH.

CONTEXT — Generative hyphae in context infrequent, colorless, thin-walled, moderately branched, 1.5–3 µm in diam; skeletal hyphae in context dominant, colorless to pale yellowish brown, thick-walled with a narrow lumen to subsolid, occasionally branched, straight, more or less regularly arranged, 3–5 µm in diam; binding hyphae in context abundant, colorless to pale yellowish brown, thick-walled with a narrow lumen to subsolid, flexuous, frequently branched, interwoven, 1–2.5 µm in diam.

TUBES — Generative hyphae infrequent, colorless, thin-walled, moderately branched, 1.2–2 µm in diam; skeletal hyphae in trama dominant, colorless to pale yellowish brown, thick-walled, occasionally branched, more or less straight, interwoven, 2.4–4 µm in diam; binding hyphae in trama colorless to pale yellowish brown, thick-walled with a narrow lumen to subsolid, flexuous, frequently branched, interwoven, 1–2 µm in diam. Cystidia absent, while cystidioles present, fusoid to tubular, sometimes tips of cystidioles branched and septate, colorless, thin-walled, 22–34 × 2.5–4.5 µm; basidia clavate, bearing four sterigmata and a basal clamp connection, 18–24 × 3.5–5 µm; basidioles in shape similar to basidia, but distinctly smaller.

SPORES — Basidiospores cylindrical, colorless, thin-walled, smooth, IKI–, CB–, (3.8–)3.9–5 × (1.8–)1.9–2.1(–2.5) µm, L = 4.34 µm, W = 2.04 µm, Q = 2.13 (n = 9/1).

TYPE OF ROT — White rot.

ADDITIONAL SPECIMENS EXAMINED — **CHINA. FUJIAN PROVINCE**, Wuyi Mountains, Longfenggu Forest Park, on fallen angiosperm trunk, 27 Aug 2006, B.K. Cui 4124 (BJFC 485, IFP 1196); Wuyishan Nature Reserve, Taoyuanyu, on fallen angiosperm trunk, 24 Aug 2006, Y.C. Dai 7376 (IFP 11887); **HUNAN PROVINCE, SHIMEN COUNTY**, Hupingshan Nature Reserve, on fallen angiosperm trunk, 16 Sep 2009, Y.C. Dai 11360 (BJFC 7283); **JIANGXI PROVINCE, FENYI COUNTY**, Dagang Mountain, on fallen trunk of *Cyclobalanopsis blakei*, 18 Sep 2008, Y.C. Dai 10420 (BJFC 4669).

Neofomitella rhodophaea (Lév.) Y.C. Dai, Hai J. Li & Vlasák, **comb. nov.** FIGS 2d,e

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≡ *Polyporus rhodophaeus* Lév., Ann. Sci. Nat., Bot., 3e Sér., 2: 190 (1844).

≡ *Fomitella rhodophaea* (Lév.) T. Hatt., Mycoscience 46: 305 (2005).

SPECIMENS EXAMINED — JAPAN. KAGOSHIMA PREF., KIMOTSUKI-GUN, Uchindura, Oct 1962, K. Aoshima (BJFC: ex. O 10866); TOKYO, Meguro, 15 Oct 1943 (BJFC: ex. TFM F-10861); MALAYSIA. NEGERI SEMBILAN, Pasoh, Sembilan For. Nat. Res., 25 Dec 1997, T. Hattori (BJFC: ex. TFM 18313).

OTHER SPECIMENS EXAMINED — *Fomitella supina* (FIG. 2f): BRAZIL. ALAGOAS STATE, PILAR MUNICIPALITY, Reserva Particular do patrimonio Natural (PPPN) São Pedro, on dead hardwood, Oct 2000, B. Tatiana, N. Gibertoni 357 (BJFC: ex. O 10849). COLOMBIA. DEPT. DE ANTIOQUIA, MUNICIPIO CALDAS FINCA, alt. 1700 m, 25 Jun 1978, L. Ryvarden 16 590 (BJFC: ex. O 10769). GUATEMALA. Lago Atitlán, on hard wood, J. Vlasák JV0610. PANAMA. Ensenada de Santa Cruz. Parque Nac. de Coiba, 17 Nov 1996, M. Nunez 1183 (BJFC: ex. O 10770). PUERTO RICO. Toro Negro, Commonwealth For., on deciduous wood, 24 Jun 1996, L. Ryvarden 39 027 (BJFC: ex. O 10772).

Discussion

The new genus *Neofomitella* is composed of two distinct lineages, one comprising *Neofomitella rhodophaea* and *N. fumosipora* and the other *N. polyzonata*, newly described from China. The main morphological characters of *Fomitella supina* and the three *Neofomitella* species are presented in TABLE 2. Like *Neofomitella polyzonata*, *N. fumosipora* has more or less brown basidiocarps; however, it has a glabrous pileal surface and smaller pores (7–9 per mm; Hattori 2005). All three *Neofomitella* species show distinctly crusted basidiocarps with a cuticle that develops from base to margin (FIG. 2a–e). In *F. supina* the cuticle also develops from the base but does not usually extend to the very margin (FIG. 2f).

Neofomitella is phylogenetically close to several *Microporus* species (FIG. 1). Although *Microporus* also has a trimitic hyphal system, colorless, thin-walled, smooth, non-dextrinoid, inamyloid basidiospores, its species usually produce stipitate basidiocarps with a white to cream context (Gilbertson & Ryvarden 1986; Núñez & Ryvarden 2001). Its type species, *M. perula* P. Beauv. [= *M. xanthopus* (Fr.) Kuntze], also has an encrusted pileal surface, but its centrally or laterally stipitate and usually infundibuliform basidiocarps, distinctly small pores, and white context (Núñez & Ryvarden 2001) clearly differentiate it from *Neofomitella* species.

Morphologically, *Corioloopsis* shares many features with *Fomitella* and *Neofomitella*, such as a more or less brown context, trimitic hyphal system with clamped generative hyphae, colorless basidiospores, and causing a white rot (Gilbertson & Ryvarden 1986; Núñez & Ryvarden 2001; Hattori 2005). However, phylogenetic analysis shows *Corioloopsis* as polyphyletic with its type species, *Polyporus occidentalis* Klotzsch [= *Corioloopsis occidentalis* (Klotzsch) Murrill; = *Trametes polyzona* (Pers.) Justo], clustering within the *Trametes* clade and distinctly separated from *Fomitella* and *Neofomitella* (FIG. 1).

TABLE 2. Main morphological characters of *Fomitella* and *Neofomitella* species.

SPECIES	BASIDIOCARPS ¹	PORES (/mm)	BASIDIOSPORES ² (µm)	CYSTIDIOLES ³	DISTRIBUTION ⁴
<i>F. supina</i>	A/P, V then G	5–7	6.5–9 × 2.4–3.5, C	–	T/S, America, Africa
<i>N. fumosipora</i>	A/P, G	6–10	3–4 × 1.7–2.2, C/OE	+	T/S, Asia
<i>N. polyzonata</i>	A, V	3–4	3.9–5 × 1.9–2.1, C	+	S/WT, China
<i>N. rhodophaea</i>	A/P, G	7–8	3.5–4.5 × 2.5–3, OE	–	T, also S/WT, Asia, Africa

¹ A = annual, P = perennial, G = glabrous, V = velutinate; ² C = cylindrical, OE = oblong-ellipsoid;

³ + = presence, – = absence; ⁴ T = tropics, S = subtropics, WT = warm-temperate.

The transfer of this taxon to *Trametes* was suggested by Corner (1989) and validated by Justo & Hibbett (2011). *Coriolopsis strumosa* (Fr.) Ryvarden usually has encrusted basidiocarps that are similar to *Neofomitella*, but *C. strumosa* has a soft corky, olivaceous-brown, umber or hazel-brown context and distinctly larger basidiospores (7–10 × 3–4 µm; Núñez & Ryvarden 2001; Li 2013).

Funalia species cluster in the same clade with *Fomitella*. Both genera share trimitic hyphal system and more or less similar basidiospores, but *Funalia* species usually produce a strongly tomentose to hispid pileal surface, a white, cream to straw colored context, and cyanophilous skeletal hyphae (Niemelä et al. 1992; Dai 1996).

Hexagonia has pileate brown basidiocarps, tissues that darken in KOH, a trimitic hyphal system, and colorless thin-walled basidiospores similar to those in *Fomitella* and *Neofomitella* (Núñez & Ryvarden 2001). However, *Hexagonia* species usually have larger hexagonal pores and distinctly larger basidiospores (usually longer than 10 µm; Gilbertson & Ryvarden 1986; Núñez & Ryvarden 2001).

No sequences of *Fomitella malaysiana* are available at present. The distinctly encrusted pileal surface (Hattori & Sotome 2013) indicates that this species may belong to *Neofomitella*. Further molecular studies are needed to resolve its taxonomic and phylogenetic position.

Key to species of *Neofomitella*

1. Pores 3–4 per mm *N. polyzonata*
1. Pores 6–10 per mm 2
2. Basidiospores 1.7–2.2 µm wide *N. fumosipora*
2. Basidiospores 2.5–3 µm wide *N. rhodophaea*

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